

STIC Search Report Biotech-Chem Library

STIC Database Technology

TO: Marcela Cordero Garcia

Location: REM-3C18 Monday, April 18, 2005

Art Unit: 1654

Spare Notes

Case Serial Number: 10/659179

From: David Schreiber

Location: Biotech-Chem Library

Remsen E01A61

Phone: 571-272-2526

David.Schreiber@uspto.gov

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STIC-Biotech/ChemLib

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From: Sent: To: Subject:	Unknown@Unknown.com Tuesday, March 29, 2005 2:59 PM STIC-Biotech/ChemLib Generic form response	·
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MyDate=Tue Mar 29 14	:58:00 EST 2005	
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Name=Marcela M Corde:	ro Garcia	
Empno=80381		
Phone=2-2939		
Artunit=1654		~ ,
Office=REM3C18		N.
Serialnum=10/659,179		:
PatClass=514/2		
Earliest=9/9/03		
Searchtopic=Please se	earch in NPL and MARPAT:	,
where: Cbz= benzyloxycarbon; Mpg=3-methoxypropylg; (if too many hits, p. 2) IF only Applicant':	lycine (a hydrophobic unnatural amino lease use "hemicalcium salt" instead constant of the search broad sown work found, please search broad	of multivalent salt) claim:
	utically acceptable multivalent metal n having a neutral thrombin S1-binding ng moiety.	
Thanks,		
Marcela		
**************************************	**************************************	****************************** Vendors and cost where applicable
Searcher: 1.5chve.bev Searcher Phone: 2-2524 Date Searcher Picked up: 41 Date Completed: 41 Searcher Prep/Rev. Time: 17 Online Time: 7	NA#: AA#: Interference: SPDI: S/L: Oligomer: Encode/Transl: Structure#: / Text: Inventor: Litigation:	STN: 3 4 4 COST WHERE APPROXIMATION OF THE PROPERTY OF THE PRO

Corrents=Please also do an inventor search:
DEADMAN, JOHN JOSEPH; MADGE, DAVID JONATHAN; DOLMAN, MARK; KAKKAR, SANJAY KUMAR; KENNEDY,
ANTHONY JAMES; COMBE-MARZELLE, SOPHIE MARIE;
CHAHWALA, SURESH BABUBHAI; BOUCHER, OLIVER VIMPANY ARNOLD

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STAFF USE ONLY	Type of Search	Vendors and cost where applicable				
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Searcher:	NA#: AA#:	DIALOG:				
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Searcher Prep/Rev. Time:	Structure#: Text:	WWW/Internet:				
Online Time:	Inventor: Litigation:	Other(Specify):				

=> fil hcaplus
FILE 'HCAPLUS' ENTERED AT 09:05:38 ON 18 APR 2005
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NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 38

STEREO ATTRIBUTES: NONE

L23 9 SEA FILE=REGISTRY SSS FUL L21

L24 3 SEA FILE=HCAPLUS L23

=> d ibib abs hitstr 124 1-3

L24 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:198296 HCAPLUS

DOCUMENT NUMBER: 140:229444

TITLE: Boronic acid salts and use thereof in the preparation

of medicaments for treating thrombosis

INVENTOR(S): Madge, David Jonathan; Dolman, Mark; Combe-Marzelle,

Sophie Marie; Deadman, John Joseph; Kennedy, Anthony

James; Kakkar, Sanjay Kumar

PATENT ASSIGNEE(S): Trigen Limited, UK

SOURCE: Eur. Pat. Appl., 19 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA'	CENT	NO.			KIND DATE			APPLICATION NO.					DATE				
EP	1396	270			A1		2004	0310	EP 2003-255629					20030909			
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		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	ΕE,	HU,	SK	
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		•	•	•	•	•	CM,	•	•		•	•	•	•	•	•	
WO	2004						2004			WO 2003-GB3887 BA, BB, BG, BR, BY, B					0030		
	W:																
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	1744 -	•	•	•		•	TM,			•		•		-		-	•
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WO	2004	•	•	•	•	•	2004	•	GN, GQ, GW, ML, MR, NE, WO 2003-GB3897					•			
5	W:		_				AU,								_		
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                           Α1
                                                                      20030909
                                             US 2003-659179
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                           A1
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PRIORITY APPLN. INFO.:
                                              GB 2002-20764
                                                                  A 20020909
                                              GB 2002-20822
                                                                  A 20020909
                                              GB 2003-7817
                                                                  A 20030404
                                              GB 2003-11237
                                                                  Α
                                                                     20030516
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                                                                      20030704
                                              US 2003-485786P
                                                                  Ρ
                                                                      20030708
                                              EP 2003-255590
                                                                  A3 20030909
OTHER SOURCE(S):
                          MARPAT 140:229444
     Salts of a peptide boronic acid drug, for example of Cbz-(R)-Phe-(S)-Pro-
     (R)-Mpg-B(OH)2 are described. The counter-ion to the boronate may be an
     alkali metal or derived from an organic nitrogen-containing compound The
salts are
     used for the manufacture of a medicament for treating thrombosis.
TΤ
     667917-16-0P, TRI 50c
     RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); PRP
     (Properties); RCT (Reactant); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
     (Reactant or reagent); USES (Uses)
        (preparation, antithrombotic activity, bioavailability and properties of
        oral boronic acid salts)
```

Absolute stereochemistry.

RN

CN

667917-16-0 HCAPLUS

4-methoxybutyl]- (9CI) (CA INDEX NAME)

L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-

IT 667917-16-0DP, complexes with tri 50c 667917-80-8P 667917-82-0P 667917-83-1P 667917-86-4P 667917-88-6P 667917-90-0P

RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation, antithrombotic activity, bioavailability and properties of oral boronic acid salts)

RN 667917-16-0 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 667917-80-8 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]-, lithium salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

•x Li

RN 667917-82-0 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]-, sodium salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

●x Na

RN 667917-83-1 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]-, potassium salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

●x K

RN 667917-86-4 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]-, compd. with L-arginine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 667917-16-0

CMF C27 H36 B N3 O7

Absolute stereochemistry.

CM 2

CRN 74-79-3 CMF C6 H14 N4 O2

'Absolute stereochemistry.

RN 667917-88-6 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]-, compd. with L-lysine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 667917-16-0 CMF C27 H36 B N3 O7

Absolute stereochemistry.

CM 2

CRN 56-87-1 CMF C6 H14 N2 O2

Absolute stereochemistry.

RN 667917-90-0 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]-, compd. with 2-deoxy-2-(methylamino)-D-glucose (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 667917-16-0 CMF C27 H36 B N3 O7

Absolute stereochemistry.

CM 2

CRN 3329-30-4 CMF C7 H15 N O5

Absolute stereochemistry.

REFERENCE COUNT:

9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L24 ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:198295 HCAPLUS

DOCUMENT NUMBER:

140:229443

TITLE:

INVENTOR(S):

Boronic acid salts of multivalent metals used in the preparation of a medicament for treating thrombosis Madge, David Jonathan; Dolman, Mark; Combe-Marzelle, Sophie Marie; Deadman, John Joseph; Kennedy, Antony James; Kakkar, Sanjay Kumar; Chahwala, Suresh Babubhai; Boucher, Oliver Vimpany Arnold; Walter, Armin; Olbrich, Alfred; Krimmer, Dieter;

Weiland-Weibel, Andrea Maria Theresia PATENT ASSIGNEE(S): Trigen Limited, UK

SOURCE:

Eur. Pat. Appl., 12 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PAT	FENT	NO.			KIN	D	DATE APPLICATION NO.					DATE						
EP	1396	 269			A1	_	20040310 EP 2003-255604				20030909							
	R:		BE.	CH.		DK.	ES,							NT.				
		IE,	SI,				RO,										11,	
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EP 1466916 A1 20041013 EP 2004-76510 20030909 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK EP 1466917 20041013 A1 EP 2004-76521 20030909 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK PRIORITY APPLN. INFO.: GB 2002-20764 A 20020909 GB 2002-20822 Α 20020909 GB 2003-7817 A 20030404 GB 2003-11237 A 20030516 GB 2003-15691 A 20030704 US 2003-485786P Ρ 20030708 EP 2003-255590 A3 20030909

OTHER SOURCE(S): MARPAT 140:229443

AB Salts of a pharmaceutically acceptable divalent metal and an organoboronic acid as selective thrombin inhibitors are described. Examples of such metals are calcium, magnesium and zinc. The organoboronic acid drug may be a boropeptide protease inhibitor. The salts may be formulated in oral dosage form, such as a capsule or compressed tablet.

IT 667917-16-0P, TRI 50C

RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation, antithrombotic activity, bioavailability and properties of oral boronic acid salts of multivalent metals)

RN 667917-16-0 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 667917-15-9P 667917-16-0DP, Complexes with zinc or magnesium

RL: PAC (Pharmacological activity); PKT (Pharmacokinetics); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation, antithrombotic activity, bioavailability and properties of oral boronic acid salts of multivalent metals)

RN 667917-15-9 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]-, calcium salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

●x Ca

RN 667917-16-0 HCAPLUS

CN L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[(1S)-1-borono-4-methoxybutyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L24 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1995:508296 HCAPLUS

DOCUMENT NUMBER: 122:281427

TITLE: Characterization of a Class of Peptide Boronates with

Neutral P1 Side Chains as Highly Selective Inhibitors

of Thrombin

AUTHOR(S): Deadman, John J.; Elgendy, Said; Goodwin, Christopher A.; Green, Donovan; Baban, Jehan A.; Patel, Geeta;

A.; Green, Donovan; Baban, Jehan A.; Patel, Geeta; Skordalakes, Emmanuel; Chino, Naoyoshi; Claeson,

Goran; et al.

CORPORATE SOURCE: Thrombosis Research Institute, London, SM2 5TF, UK

SOURCE: Journal of Medicinal Chemistry (1995), 38(9), 1511-22

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society

DOCUMENT TYPE:

American chemical society

Journal

LANGUAGE: English

GΙ

AΒ Z-D-Phe-Pro-boroMpg-OPin (I) has been shown previously to be a highly specific inhibitor of thrombin in spite of lacking an arginine-like guanidino group at the Pl site. A range of compds. have been synthesized based upon this lead compound, varying the neutral side chain at the P1 site. Of the 20 examples based upon the structures at P2 and P3 of Z-D-X-Pro (X being Phe or β , β -diphenylalanine), all were effective inhibitors of thrombin (Ki's between 10 and 100 nM). Furthermore all exhibited a high specificity toward thrombin having values for a Ki(trypsin)/Ki(thrombin) ratio of between 10- and 100-fold. High ratio values were found for a number of the compds. tested against a range of serine proteinases (plasmin, factor Xa, kallikrein, urokinase, protein Ca, chymotrypsin, elastase, and cathepsin G).2. As far as potency toward thrombin, compds. containing the methoxypropyl group at P1 were favored over those with a methoxy grouping on a shorter alkyl chain (8) or without the methoxy group (1-5). The compds. display potent anticoagulant activity with values for 18 in thrombin time of 0.63 μM and in activated partial thromboplastin time of 2.0 μM . 11B NMR has been used to confirm interaction of the boron atom with the active site. From the high specificity shown with all the compds., the authors propose that the compds., constitute a new class of thrombin inhibitors.

IT 162854-83-3P

CN

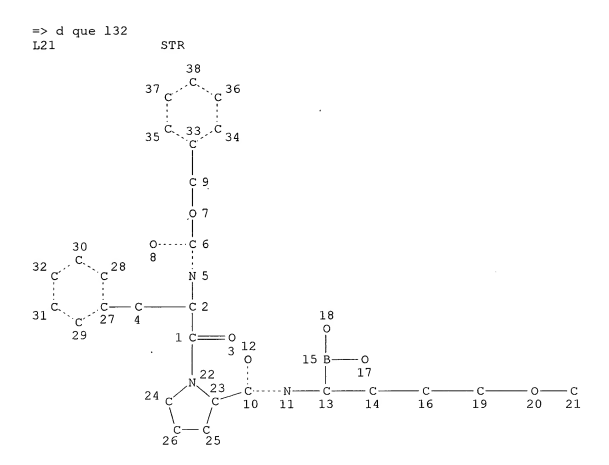
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(characterization of a class of peptide boronates with neutral P1 side chains as highly selective inhibitors of thrombin in relation to anticoagulant activity)

RN 162854-83-3 HCAPLUS

L-Prolinamide, N-[(phenylmethoxy)carbonyl]-D-phenylalanyl-N-[1-(dimethoxyboryl)-4-methoxybutyl]-, (R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 38

STEREO ATTRIBUTES: NONE

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L25	9551	SEA	FILE=HCAPLUS	INHIBIT	?(5A)THROMBIN?
L26	81	SEA	FILE=HCAPLUS	L25 AND	(BORON? OR ORGANOBORON?)
L27	6	SEA	FILE=HCAPLUS	L26 AND	S1?
L28	. 3	SEA	FILE=HCAPLUS	L26 AND	S2?
L29	3	SEA	FILE=HCAPLUS	L26 AND	S3?
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=> d ibib abs 132 1-8

L32 ANSWER 1 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 2003:186947 HCAPLUS

TITLE: Thrombin

AUTHOR(S):

CORPORATE SOURCE:

SOURCE:

Rohde, Rosemary; Maderna, Andreas; Hawthorne, Fred UCLA, Dept. of Chemistry, Los Angeles, CA, 90095, USA Abstracts of Papers, 225th ACS National Meeting, New Orleans, LA, United States, March 23-27, 2003 (2003), PRES-074. American Chemical Society: Washington, D.

C.

CODEN: 69DSA4

DOCUMENT TYPE: Conference; Meeting Abstract

LANGUAGE: English

Serine proteases constitute a major class of enzymes that are widely AΒ distributed in the human body. Examples of serine proteases are thrombin, trypsin, chymotrypsin, factor Xa, and human leukocyte elastase (HLE). Serine proteases are frequently the cause of many life-threatening diseases. Thrombin, for example, plays a central role in thrombosis and hemostasis. Thrombosis, or excessive blood clotting, is the major culprit of numerous cardiovascular diseases. Due to the high mortality rate of this disease and others, there is intensive interest in developing an orally active thrombin inhibitor. The goal of this work is to design and produce a selective and orally active thrombin inhibitor by synthesizing a boronated trans-lactum that will specifically bind to the active site of the enzyme. The thrombin inhibitor FE-1, composed of a trans-lactam template, a piperazinyl bisamide linker, and a hydrophobic carborane cage, was designed and optimized using mol. modeling and protein-ligand docking calcns. These techniques were utilized to determine the correct size, orientation, and stability needed for the inhibitor. FE-1 was synthesized in 22 steps and shown to have very good interactions with active site residues of thrombin: the Me groups on the carborane make lipophilic contact with the benzene ring to tryptophan 215 and isoleucine 174 and the S3 pocket; the carbonyl group of the amide makes hydrogen bonds with the amino group of glycine 216; the piperazine ring has great contact with tyrosine 60A; and the trans-lactam template in its transition state is oriented in such a way that the OH group of serine 195 cleaves the amide bond of the lactam resulting in an intermediate tetrahedral carbon center in which the oxyanion group is place in the oxyanion hole of the S-1 site. All these interactions are extremely important in the development of thrombin-specific anticoagulants. In order to achieve an optically pure synthetic thrombin inhibitor, chiral HPLC was used in the final step of the reaction in order to sep. the correct a-diastereomer. This is unique and unprecedented mol. assemble represents an example of a new class of boronated enzyme inhibitors and yielded a new potential thrombin inhibitor. Partially funded by the ACS Scholars Program.

L32 ANSWER 2 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:354554 HCAPLUS

DOCUMENT NUMBER: 137:87844

TITLE: Design of Selective Thrombin

Inhibitors Based on the (R)-Phe-Pro-Arg

Sequence

AUTHOR(S): Danilewicz, John C.; Abel, Stuart M.; Brown, Alan D.;

Fish, Paul V.; Hawkeswood, Edward; Holland, Stephen J.; James, Keith; McElroy, Andrew B.; Overington,

John; Powling, Michael J.; Rance, David J.

CORPORATE SOURCE: Departments of Discovery Chemistry, Drug Metabolism,

Discovery Biology, and Molecular Informatics Structure and Design, Pfizer Global Research and Development,

Searched by David Schreiber 22526 Page 13

Sandwich, Kent, CT13 9NJ, UK

Journal of Medicinal Chemistry (2002), 45(12), SOURCE:

2432-2453

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal English LANGUAGE:

OTHER SOURCE(S): CASREACT 137:87844

Potent and selective inhibitors of thrombin were

sought based on the (R)-Phe-Pro-Arg sequence. The objective was to generate similar binding interactions to those achieved by potent competitive inhibitors of the argatroban type, so eliminating the need for covalent interaction with the catalytic serine function, as utilized by aldehyde and boronic acid type inhibitors. Improving the \$1 subsite interaction by substitution of arginine with a 4-alkoxybenzamidine residue provided potent lead 2 (Ki = 0.37 nM). an amide bond, which H-bonds to the active site, is lost, modeling indicated that a new H-bond is generated between the alkoxy oxygen atom and the catalytic Ser-195 hydroxyl group. Substitution of the benzamidine system by 1-amidinopiperidine then gave compound 4, which provided a further gain in selectivity over trypsin. However, previous work had shown that these compds. were likely to be too lipophilic (log D +0.4 and +0.2, resp.) and to suffer rapid hepatic extraction, presumably via biliary elimination. Accordingly, both proved short-acting when administered i.v. to rats and showed poor activity when given intraduodenally. The aim was then to reduce lipophilicity below a log D of -1.2, which in a previously reported series had been effective in preventing rapid clearance. It was anticipated that compds. of this type would rely on the cation selective paracellular route of absorption from the gastrointestinal tract. Potent polar analogs with selectivity >1000 over trypsin were obtained. The best in vivo activity was shown by compound 12. However, in the final anal., its oral bioavailability proved poor, relative to analogs with similar physicochem. properties derived from argatroban, consistent with the hypothesis that mol. shape is an addnl. important determinant of

paracellular absorption. REFERENCE COUNT: THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS 46 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L32 ANSWER 3 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN

2000:619068 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 134:171

TITLE: Selective boron-containing thrombin

inhibitors-X-ray analysis reveals surprising

binding mode

von Matt, A.; Ehrhardt, C.; Burkhard, P.; Metternich, AUTHOR(S):

R.; Walkinshaw, M.; Tapparelli, C.

CORPORATE SOURCE: Novartis Pharma AG, Basel, CH-4002, Switz.

SOURCE: Bioorganic & Medicinal Chemistry (2000), 8(9),

2291-2303

CODEN: BMECEP; ISSN: 0968-0896

Elsevier Science Ltd. PUBLISHER:

DOCUMENT TYPE: Journal LANGUAGE: English

Based on the structural comparison of the S1 pocket in different trypsin-like serine proteases, a series of Boc-d-trimethylsilylalanineproline-boro-X pinanediol derivs., with boro-X being different amino

boronic acids, have been synthesized as inhibitors of thrombin. Among the novel compds., a number of derivs. were

synthesized which appeared to have side-chain variants too big to fit into

the S1 pocket. Nevertheless, these compds. inhibited thrombin in the nM range. The x-ray structure of one of these inhibitors bound to the active side of thrombin reveals that a new binding mode is responsible for these surprising results.

REFERENCE COUNT: 78 THERE ARE 78 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L32 ANSWER 4 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:605764 HCAPLUS

DOCUMENT NUMBER: 129:341097

TITLE: Bifunctional Peptide Boronate

Inhibitors of Thrombin:

Crystallographic Analysis of Inhibition

Enhanced by Linkage to an Exosite 1 Binding Peptide

AUTHOR(S): Skordalakes, Emmanuel; Elgendy, Said; Goodwin,

Christopher A.; Green, Donovan; Scully, Michael F.; Kakkar, Vijay V.; Freyssinet, Jean-Marie; Dodson, Guy;

Deadman, John J.

CORPORATE SOURCE: Peptide Synthesis Section and Biochemistry Section,

Thrombosis Research Institute, London, SW3 6LR, UK

SOURCE: Biochemistry (1998), 37(41), 14420-14427

CODEN: BICHAW; ISSN: 0006-2960

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

The affinity of the hirudin49-64 segment for exosite 1 of thrombin has been used previously to enhance the potency of simple competitive inhibitors [DiMaio, J., Gibbs, B., Munn, D., Lefebvre, J., Ni, F., Konishi, Y. (1990) J. Biol. Chemical 265, 21698-21703, and Maraganore, J. M., Bourdon, P., Jablonski, J., Ramachandran, K. L., and Fenton, J. W., II (1990) Biochem. 29, 7095-7087]. Using a similar approach, we have enhanced the activity of two active site directed thrombin inhibitors by attaching this segment via a novel reverse oriented linker to each of two tripeptide boronate inhibitors. At P1, compound 1 contains an arginine-like, isothiouronium, side chain, while compound 2 contains an uncharged, bromopropyl residue. Inhibition of human α - thrombin by compound 1 shows slow, tight-binding competitive kinetics (final Ki of 2.2 pM, kl of 3.51+107 M-1 s-1, and k-1 of 1.81+10-4 s-1). The addition of hirugen peptide (20 μM) competes for exosite 1 binding and restores the k1 and k-1 to that of the analogous tripeptide, 0.29+107 M-1 s-1 and 0.13+10-4 s-1, Compound 1 has enhanced specificity for thrombin over trypsin with KiTry/KiThr of .apprx.900 compared to the analogous tripeptide, with KiTry/KiThr of .apprx.4. Compound 2 acts as a competitive inhibitor (KiThr of 0.6 nM) and is highly selective with no effect on trypsin. Crystallog. anal. of complexes of human α -thrombin with compound 1 (1.8 Å) and compound 2 (1.85 Å) shows a covalent bond between the boron of the inhibitor and Ser195 (bond lengths B-O of 1.55 and 1.61 Å, resp.). The isothiouronium group of compound 1 forms bidentate interactions with Asp189. The P2 and P3 residues of the inhibitors form interactions with the \$2 and \$3 sites of thrombin similar to other D-Phe-Pro based inhibitors [Bode, W., Turk, D., and Karshikov, A. (1992) Protein Sci. 1, 426-471.]. The linker exits the active site cleft of thrombin forming no interactions, while the binding of Hir49-64 segment to exosite 1 is similar to that previously described for hirudin [Rydel, T. J., Tulinsky, A., and Bode, W. (1991) J. Mol. Biol. 221, 583-601.]. Because of the similarity of binding at each of these sites to that of the analogous peptides added alone, this approach may be used to improve the inhibitory activity of all types of active site directed thrombin

inhibitors and may also be applicable to the design of inhibitors of other proteases.

REFERENCE COUNT:

THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L32 ANSWER 5 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN

42

ACCESSION NUMBER:

1997:671089 HCAPLUS

DOCUMENT NUMBER:

127:341584

TITLE:

Selection of \$18326 as a new potent and

selective boronic acid direct

thrombin inhibitor

AUTHOR(S):

Rupin, A.; Mennecier, P.; Lila, C.; De Nanteuil, G.;

Verbeuren, T. J.

CORPORATE SOURCE:

Div. Angiology, Servier Research Inst., Suresnes,

F-92150, Fr.

SOURCE:

Thrombosis and Haemostasis (1997), 78(4), 1221-1227

CODEN: THHADQ; ISSN: 0340-6245

PUBLISHER:

Schattauer DOCUMENT TYPE: Journal LANGUAGE: English

Using enzymic microassays, the potency of a series of new boroArg tripeptides was determined vs. thrombin and a panel of Ser proteases implicated in the coagulation and fibrinolysis pathways. The inhibition of the Ser protease complement factor I was also studied. Factor I regulates the alternate pathway of the complement and its inhibition appears to be responsible for the toxic effects of the orally available thrombin inhibitor Ac-D-Phe-Pro-boroArg (DuP-714). The structure of the new boronic acid derivs. tested was modified from that of DuP-714 by replacing the proline in the P2 position by N-cycloalkylglycine residues of increasing size (\$18989: cyclopropyl; \$18563 : cyclobutyl; **S18326**: cyclopentyl; **S18229**: cyclohexyl). All compds. were found to be slow-tight binding inhibitors of thrombin vs. purified human fibrinogen. Replacement of Pro by N-cycloalkylglycines did not decrease the anti-thrombin potency of the substances up to the cyclopentyl size and this result was confirmed by classical coagulation assays with human plasma in vitro. In contrast, the inhibitory activities of the 4 new boronic acids were found to be lower than those of DuP-714 vs. plasmin, urokinase (u-PA), plasmatic kallikrein, activated protein C (aPC) and complement factor I. The cyclopentyl derivative $\bf S18326$ is a slightly more active inhibitor of thrombin than DuP-714 (initial IC50 values 3.99 nM vs. 4.73 nM, resp.). Moreover \$18326 was identified as the most selective compound of the series with relative potencies being 2-29-fold higher than that of DuP-714 vs. the panel of Ser -proteases tested; the rank order of potency vs. the other Ser proteases for \$18326 was t-PA > kallikrein > aPC > factor I > plasmin > fXa > u-PA. These results indicate that the size of the thrombin hydrophobic pocket \$2 is sufficient to accept larger residues than Pro in the P2 position of Ac-D-Phe-X-boroArg derivs. while this is not the case for other important Ser proteases of the fibrinolysis, coagulation, and complement pathways. The N-cyclopentyl glycine containing derivative S18326, which is the most potent and the most selective anti-thrombin compound of the series, currently undergoes

L32 ANSWER 6 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

major preclin. testing.

1997:667172 HCAPLUS

DOCUMENT NUMBER:

127:244735

TITLE:

Crystallographic Structures of Human $\alpha\text{-Thrombin}$

Complexed to Peptide Boronic Acids Lacking a

Positive Charge at Pl. Evidence of Novel Interactions Skordalakes, Emmanuel; Tyrell, Richard; Elgendy, Said; Goodwin, Christopher A.; Green, Donovan; Dodson, Guy; Scully, Michael F.; Freyssinet, Jean-Marie H.; Kakkar,

Vijay V.; Deadman, John J.

CORPORATE SOURCE:

AUTHOR(S):

SOURCE:

Thrombosis Research Institute, London, SW3 6LR, UK Journal of the American Chemical Society (1997),

119(41), 9935-9936

CODEN: JACSAT; ISSN: 0002-7863

Moc-Dpa-Pro-boroMpg, compound (I), lacking a pos. charge at P1 is a potent

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

inhibitor of human α - thrombin (H α t) (KiThr = 3nM). The crystallog. anal. of the enzyme:inhibitor complex of I at 1.9Å resolution, provides for the first time a partial explanation for the basis of the high affinity interaction at the s1 site. Tripeptide boronates I and Z-Dpa-Pro-boroVal, compound (II), were synthesized as described, and crystals obtained for I and II with $H\alpha T$ and N-Ac-hirugen. Crystals were flash cooled and data sets were collected to a maximum Bragg spacing of 1.8Å and 2.1Å resp. and subsequently processed with Denzo and Scalepack and AMORE (HαT.hirugen.PPACK). The data was further refined using Spartan, Refmac and ARP. Refinement converged to a crystallog. R factor of 17.5% (Rfree = 24.0%, using 5% of reflections) and 17.0% (Rfree = 23.5%) and R-factors were 0.32 and 0.36, and RMS deviations were 0.02Å and 2.4°, and 0.019Å and 2.5° for the complex with I and II, resp. Atomic coordinates have been deposited in the Brookhaven Protein Data Bank. Both compound I and II form the canonical interactions with human α -thrombin at the S2 and S3 sites, already shown with the PPACK complex. Complex I shows the expected covalent interaction of c.a. 1.75Å between the boron and the Ser-1950γ of the $H\alpha T$ and O1B is coordinated by Gly-193NH and Ser-195NH in the so called oxy-anion pocket (Figure 1) (01B-193GlyNH 2.79Å, 01B-ser195NH 3.11Å). In complex I, the ether oxygen is functioning as a hydrogen bond acceptor from a water (2.54 Å) which is, in turn, bridging to Gly-216CO and Gly-219CO. This bridging interaction has previously been observed in the fibrinopeptide A - α - thrombin complex, between the ε-NH of the arginine guanidino, WAT80 and Gly219CO. Surprisingly, despite the reasonable affinity of compound II, (Kithr 20 nM), crystallog. anal. at 2.1 Å of the complex II shows a novel interaction where the boron is 3.34Å from Ser-1950y, and the boron oxygen OlB is now displaced from the oxyanion pocket and is hydrogen bonded (2.84Å) to Ser-1950 γ . The displacement allows O1A to interact more strongly with the carboxylate side chain of Glu-192 (O1A-Glu192OE1 3.11Å compared to 4.16Å for complex II and I

 α -thrombin may provide a better understanding for the design of low mol. weight inhibitors of high specificity. THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS 22

displaced into close proximity with Val-213 of $H\alpha T$. The discovery

resp.). The inhibitor P1 valine-like iso-Pr side chain in complex II is

REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L32 ANSWER 7 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN

of this interaction between \$1 and \$3 for human

ACCESSION NUMBER: 1997:56330 HCAPLUS

DOCUMENT NUMBER: 126:139500

TITLE: S1 heterocyclic thrombin

inhibitors

AUTHOR(S): Dominguez, C.; Carini, D. J.; Weber, P. C.; Knabb, R.

M.; Alexander, R. S.; Kettner, C. A.; Wexler, R. R.

CORPORATE SOURCE: Exptl. Sta., DuPont Merck Pharmaceutical Co.,

Wilmington, DE, 19880-0500, USA

SOURCE: Bioorganic & Medicinal Chemistry Letters (1997), 7(1),

79-84

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

AB A series of boropeptides have previously been described by Kettner et al.

to be potent thrombin inhibitors. DuP 714 is a

representative of this class of compds. with a Ki = 0.040 nM, but this inhibitor has undesirable side effects. New and selective **boronic**

acid thrombin inhibitors have been developed by

replacing the guanidine of the boroarginine side chain with various

heterocycles ranging in size and basicity.

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L32 ANSWER 8 OF 8 HCAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1995:413502 HCAPLUS

DOCUMENT NUMBER:

122:259717

TITLE:

Kinetic and Crystallographic Studies of Thrombin with

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS

Ac-(D) Phe-Pro-boroArg-OH and Its Lysine, Amidine,

Homolysine, and Ornithine Analogs

AUTHOR(S): Weber, Patricia C.; Lee, Sheng-Lian; Lewandowski,

Francis A.; Schadt, Margaret C.; Chang, Chong-Hwan;

Kettner, Charles A.

CORPORATE SOURCE:

REFERENCE COUNT:

Chemical and Physical Sciences Department, The Du Pont

Merck Pharmaceutical Company, Wilmington, DE,

19880-0228, USA

SOURCE:

Biochemistry (1995), 34(11), 3750-7

CODEN: BICHAW; ISSN: 0006-2960

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

LANGUAGE: English

AB The x-ray crystallog. structure of Ac-(D)Phe-Pro-boroArg-OH (DuP714, Ki = 0.04 nM) complexed with human α -thrombin shows the **boron**

0.04 nM) complexed with human α -thrombin shows the **boron** atom covalently bonded to the side-chain oxygen of the active site serine,

Ser195. The boron adopts a nearly tetrahedral geometry, and the boronic acid forms a set of interactions with the protein that

mimic the tetrahedral transition state of serine proteases. Contributions of the arginine side chain to inhibitor affinity were examined by synthesis of the ornithine, lysine, homolysine, and amidine analogs of DuP714. The basic groups interact with backbone carbonyl groups, water mols., and an aspartic acid side chain (Asp189) located in the thrombin **S1**

aspartic acid side chain (Asp189) located in the thrombin **S1** specificity pocket. The variation in inhibition constant by 3 orders of magnitude appears to reflect differences in the energetics of interactions made with thrombin and differences in ligand flexibility in solution Kinetic

and crystallog. data are reported for the following thrombin inhibitors: DuP714 (space group C2, a = 70.8 Å, b = 72.3

Å, c = 72.6 Å, $\beta = 100.6$ °, crystallog. R-factor =

0.204 to 1.95 Å resolution); Ac-(D)Phe-Pro-boroLys-OH (Ki = 0.24 nM, C2,

 $a = 70.3 \text{ Å}, b = 71.9 \text{ Å}, c = 71.9 \text{ Å}, \beta = 100.9^{\circ},$

R-factor = 0.201 to 2.35 Å resolution); Ac-(D)Phe-Pro-boro-homoLys-OH (Ki = 8.1 nM, C2, a = 70.3 Å, b = 71.9 Å, c = 71.9 Å, β =

100.9°, R-factor = 0.212 to 2.4 Å resolution);

Ac-(D)Phe-Pro-boroOrn-OH (Ki = 79 nM, C2, a = 70.4 Å, b = 72.2 Å, c = 72.2 Å, β = 100.1°, R-factor = 0.195 to 2.25 Å resolution); and Ac-(D)Phe-Pro-boro-n-butylamidinoGly-OH (Ki = 0.29 nM, C2, a = 70.8 Å, b = 72.4 Å, c = 72.2 Å, β = 100.3°, R-factor = 0.197 to 2.25 Å resolution).